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Halomethyl-1,3,5-triazines containing a photoinitiator molety.

This invention relates to photosensitive compounds, more particularly, derivatives of halomethyl-1,3,5triazines.

Halomethyl-1,3,5-triazines are known to be initiators for a number of photochemical reactions. Mixed initiators have been disclosed for use with polymerizable monomers. It would be desirable to combine a halomethyl-1,3,5-triazine with a second photoinitiator in the same molecule to provide an improved photoinitiator. This invention provides radiation-sensitive organo-halogen compounds having a photo-labile halomethyl-1,3,5-triazine molety and at least one photoinitator molety within one molecule. The compounds of this invention reare good photoinitiators, and compositions containing them can be used in printing, duplicating, copying, and maying systems. The compounds of this invent \(\mathbb{N} \) in a photocurable or photopolymerizable composition. Other imaging systems. The compounds of this invention eliminate the need for a combination of photoinitiators

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HALOMETHYL-1,3,5-TRIAZINES CONTAINING A PHOTOINITIATOR MOIETY

BACKGROUND OF THE INVENTION

Field of the invention

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This invention relates to photosensitive compounds, more particularly, derivatives of halomethyl-1,3.5-triazines.

Discussion of the Prior Art

Compounds that decompose to generate free radicals (free radical generating agents) upon exposure to
light are well known in the graphic arts. Organic halogen compounds, which are capable of generating free
radicals such as a chlorine free radical or a bromine free radical upon exposure to light, have been widely
used as photohilutors in photopolymerizable compositions, as photoactivators in free radical photographic
compositions, and as photonitilators for reactions catalyzed by adds formed by light. The spectral
compositions, and as photonitilators for reactions catalyzed by adds formed by light. The spectral
transfer tier absorbed energy to the organic halogen compound. The use of such halogen compounds in
photopolymerization processes and free radical photographic processes has been described in Kosar, LightSensitive Systems, J. Villey & Sons (New York, 1955), pp. 180-161, 343-730.

Halometry-1-3.5-triazines are known to be initiators for a number of photochemical reactions. They are employed to produce free radicals for initiating polymerization or color changes and for initiating secondary reactions upon liberation of acid by the interaction of the free-adicals when hydrogen donors are present.

Examples of the use of the halomethyl-1,3,5-triazines in the free radical polymerization of acrylate monomers are described in U.S. Patent No. 3,095,815; U.S. Patent No. 3,017,288; U.S. Patent No. 4,316,175; U.S. Patent No. 4,316,175; U.S. Patent No. 4,476,215; and DE 3,517,440; U.S. Patent No. 3,779,778 discloses the photoinitiated acid catalyzed decomposition of pyranyl ether derivatives to produce ophotosolubilizable compositions useful as positive printing plates. Chromophore substituted stryl-1,3,5-triazines and their uses are disclosed in U.S. Patent No. 3,987,037 and U.S. Patent No. 3,554,475.

Radiation sensitive compositions containing bi-and polyaromatic substituted triazines are disclosed in U.S. Patent No. 4,189,323.

Typical photopolymerization initiators for polymerizable ethylenically unsaturated compounds include 35 beazil, benzoin, benzoin ethyl ether, Michler's kethone, anthracquinone, acridine, phenazine, benzopherone, etc. For example, the use of photoinitiators such as benzoin ethers are described in U.S. Patent No. 2,722,512; anthracquinones are described in U.S. Patent No. 3,046,127; amino phenyl kethores and active ethylene or amino compounds are described in U.S. Patent No. 3,661,588; and Michler's ketone and benzophenone are described in U.S. Patent No. 3,682,641.

Mixed photoinitiators have been disclosed for use with polymerizable monomers. A very effective mixed initiator system has been Michler's ketone admixed with benzophenone, as described in U.S. Patent No. 1882.641. 3-Keto-coursains have been shown to be effective mixed initiators with Michler's ketone, as described in U.S. Patent No. 4.966.228. Halogenated organic photoinitiators such as carbon barbaromide, etc. have been used as co-photoinitiators with diazontum sates, as described in U.S. Patent No. 4.113.497. 45 Likewise, 5-isoxazolones are effective co-photoinitators when used with aromatic carbonyl compounds in the presence of halomethyl-1,3-fritains, as described in U.S. Patent No. 4.254.432. Other examples of specific combinations of known photoinitiators are disclosed in U.S. Patent Nos. 3.314,607; 3,673.140; 3.382,7(1); 3,477.11,3.47.116; 3,471.54.44 4.264.700

SUMMARY OF THE INVENTION

This invention provides radiation-sensitive organo-halogen compounds having good sensitivity in the ultraviolet and visible range of the spectrum. These compounds are suitable for use in radiation-sensitive

compositions. This invention provides compounds that have a photo-labile hatomethyl-1,35-friazine moiety and an additional photoinitiator moiety within one molecule so as to eliminate the need for a commonate of this invention are good photoinitators. Photopolymenzable and photocrosslinkable compositions containing these photoinitiators can be used in printing, duplicating, copyring, and other imaging systems.

This invention provides compounds having a 1.3.5-triazine moiety having at least one hadomethy!

This invention invoides compounds having a 1.3.5-triazine moiety having at least one hadomethy disputs and at least one additional photoinitiator moiety attached to another carbon atom of the triazine nucleus that is capable of interesting free radical or ionic chain polymerization upon exposure to actrice radiation. Representative or examples of such photoinitiator moieties include benzoin group, benzoin alkyl ether group, acatophenone group, disalkovyacetophenone group, benzophenone group, acatophenone group, disalkovyacetophenone group, benzophenone group, acatophenone group, disalkovyacetophenone group, disalkovyacetophenone group, benzophenone group, acatophenone group, disalkovyacetophenone group, d

These compounds are useful as photoinitiators for photosensitive compositions and elements. Thus, they can be incorporated in photopolymerizable compositions and printing compositions useful for producing printing plates, such as lithographic plates, relief plates or gravure plates, photoresists and photographic are comments, and photosensitive resist forming compositions with which visible images can be obtained upon exposure to light.

DETAILED DESCRIPTION OF THE INVENTION

As used herein, the term "photoinitiator moiety" means a moiety containing at least one group that is capable of initiating free radical or ionic chain polymerization upon exposure to actinic radiation.

Halomethyl-1.3.5-triazine compounds of this invention can be represented by the general formula I:

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A represents a member selected from the group consisting of mono-, di- and trihalomethyl groups, Y represents a member selected from the group consisting of A, L-P, NH₂, NHR, NH₂, OR, and R'.

Y represents a memoer securation in a goop consisting of a consistency of the carbon atoms, or a substituted or unsubstituted any group, preferable having 1 to 6 carbon atoms, or a substituted or unsubstituted any group, preferably having 1 to 6 carbon atoms, a substituted or unsubstituted any group, preferably having 1 to 6 carbon atoms, a substituted or unsubstituted any group, a substituted or unsubstituted any group, preferably having 2 to 6 carbon atoms, or a substituted or unsubstituted or unsubstituted.

heterocyclic aromatic group,
P represents a photoinitator moiety that is capable of initiating free radical or ionic chain polymerization
upon exposure to actinic radiation, and

L represents a group or a covalent bond linking the photoinitiator molety to the triazine nucleus.

Halomethyl groups that are suitable for the present invention include chloro-, bromo-, and indomethyl groups, with chloro- and bromomethyl groups being preferred. Trihalomethyl groups are preferred are inchloromethyl and tribromomethyl groups.

Y represents any of a variety of substituents that are useful in modifying the physical, e.g., solublity, or chemical properties of the molecule, and preferably represents A. L.P., or R. When Y represents A. the maximum number of habmethyl groups per triazine nucleus can be made available for free radical ceneration. When Y represents L.P. the chemical composition for both L.P groups can be the same, or it

can be different, depending on the composition of linking group L, photoinitiator moiety P, or both. When Y represents R', and in particular when R' represents an anyl, aralkenyl, or heterocyclic aromatic group, the spectral sensitivity of the molecule can be varied, based on the photochemical response of R' to actinic radiation.

When R or R' represents an aryl group it is preferred that the group have a maximum or five rings, more preferably three rings, and most preferably one ring.

When R or R' represents a substituted group, the particular identity of the substituents is not critical. However, the substituents should be selected so as not to adversely affect the photonitiation characteristics or light sensitivity of the compounds of this invention.

P preferably represents at least one group selected from the group consisting of benzoin group, benzoin alkyl ether group, an accephanene group, dialkosyacetophenone group, benzophenone group, arthraquinone group, thiosanthone group, transpullonium group, dialyridodnium group, azide group, diazonium group, 3-ketocoumarin group, bisimidazole group, fluorenone group, or a halomethyl-1,3-triazine group covalently bonded to the triazine nucleus of formula I. There is no upper triazine nucleu per photoinitiator moieles per triazine nucleus; there is no upper limit on the number of triazine nucleus. Preferably, the number of photoinitiator moiety per each triazine nucleus ranges from one to two or two to one; more preferably, there is one ephotoinitiator moiety per each triazine nucleus. If more than one photoinitiator moiety is present per triazine nucleus. We can be from different generic classes or can so be different species from the same generic class. If more that one triazine nucleus is present per hobotinitiator moiety, they can be of different species.

L represents a group that links the photoinistant moiety or molelies to the triazine nucleus. The precise identity of L is not critical, but it should be selected so that it does not interfere with or adversely affect the photoinitation characteristics or light sensitivity of the compound. L can be formed from a simple group or it are not be formed from a combination of groups. In addition, L also includes a covalent bond. Groups that are suitable for linking groups include carbamato (NHCO₂), use (NHCOMH), amino (NH), amido (CONH), aliphatic, e.g., having up to 10 carbon atoms, alsoly, e.g., having up to 10 carbon atoms, aryl, e.g., having one ring.

30 groups for attachment directly to the triazine nucleus are carbamato, urea, artino, alternyl, and other.

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One method of preparing the compounds of this invention is by the addition reaction of isocyanatosubstituted halomethyl-1,3,5-tritazines with photoinitiators having groups reactive with the isocyanate group. The isocyanato substituted tritazines may be prepared from the corresponding amino derivative according to the procedure of U. Von Gizycki, Angew. Chem. Int. Ed. Eng., 1971, 10, 403. Isocyanato-1,3,5-triazines suitable for this reaction include.

2.4-bis(trichloromethyl)-6-isocyanato-1,3,5-triazine

2-isocyanato-4-methyl-6-trichloromethyl-1,3,5-triazine

2-isocyanato-4-phenyl-6-trichloromethyl-1,3,5-triazine

2-isocyanato-4-methoxy-6-trichloromethyl-1,3,5-triazine

2-isocyanato-4-(p-methoxyphenyl)-6-trichloromethyl 1,3,5-triazine 2-isocyanato-4-(p-methoxystyryl)-6-trichloromethyl-1,3,5-triazine

2-isocvanato-4-(m.o-dimethoxyphenyl)-6-trichloromethyl-1,3,5-triazine

Typical photoinitistors that will combine with the isocyaneto group include 1-barazyl cyclohexanol (Irgacure⁶ 184), 4-hydroxyseatophenone, 4-hydroxybenzophenone, 4-minobenzophenone, 2-amino-Bituorenone, 2-amino-Bituorenone, 2-mino-Bituorenone, 2-mino-Bituorenone,

The isocyanate addition reaction can be carried out in the presence of solvents such as, for example, toluene, pyridine, benzene, xylene, dioxare, tetrahydroturan, etc., and mixtures of solvents. The duration and temperature of the reaction is dependent on the particular compounds and the catalyst employed. Generally, temperatures ranging from about 25°C to 150°C for from one to seventy-two hours are sufficient to provide for the reaction. Preferably, the reaction is carried out at room temperature from three to seventy-

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two hours. The preferred catalyst is di-n-butyltin dilaurate.

Another method for preparing the compounds of this invention is the cotrimerization of organic nitriles having a photoinitiator substituent with haloacetonitriles in accordance with the teachings of Wakabayashi et al, Bulletin of the Chemical Society of Japan, 1969, 42, 2924-30. Still another method is the condensation 5 reaction of an aldehyde compound having a photoinitiator functionality in accordance with the teachings of U.S. Patent No. 3,987,037. Another method is the nucleophilic displacement reactions on halomethyl-1,3.5triazines using photoinitiators having free hydroxy or amino groups.

The photoinitiators of this invention combine a halomethyl-1,3,5,-triazine moiety and another photoinitiator moiety in the same molecule, thereby eliminating the requirement of adding each type of photoinitiator 10 separately. Synergism can be brought about in instances where the radicals produced by one molety induce chain decomposition of the other moiety, thereby in effect, promoting the same reaction that would have been produced by a direct photoresponse.

Combinations of moieties can be selected that produce, upon exposure to actinic radiation, several different photoproducts, which will initiate the same reaction. For example, a halomethyl-1,3,5,-triazine moiety combined with a benzoin group can initiate free radical polymerization by generating halogen radicals and benzoyl radicals (along with other unidentified radical species).

Combinations of moieties can be selected that produce photoproducts which will initiate a multiplicity of reactions. For example, compounds derived from a halomethyl-1,3,5,-triazine and a sulfonlum salt can produce both free radicals and Lewis acids to cure a mixture comprising acrylate and epoxy monomers.

Furthermore, the chromophore of each photoinitator moiety can be selected so that each moiety can either (a) broadly respond to radiation, thereby allowing more efficient utilization of various light sources commercially available, or (b) narrowly respond to different wavelengths, thereby allowing stepwise initiation of multiple reactions. An example of the latter scheme would be to first irradiate the compound at a wavelength at which the halomethyl-1,3.5-triazine molety absorbs light to initiate a free radical polymeriza-25 tion, and then subsequently irradiate the compound at a wavelength at which the other photoinitiator molety absorbs light to initiate an ionic cross-linking reaction.

The sensitivity of compositions containing the compounds of this invention to actinic radiation of a particular range of wavelengths can be increased by the incorporation of ultraviolet and visible light sensitizers, such as, for example, cyanine, carbocyanine, merocyanine, styryl, acridine, polycyclic aromatic 30 hydrocarbons, polyarylamines and amino-substituted chalcones. Suitable cyanine dyes are described in U.S. Patent No. 3,495,987. Suitable styryl dyes and polyarylamines are described in Kosar. Light Sensitive Systems, J. Wiley and Sons (New York, 1965), pp 361-369. Polycyclic aromatic hydrocarbons useful as sensitizers, e.g. 2-ethyl-9,10-dimethoxyanthracene, are disclosed in U.S. Patent No. 3,640,718. Amino substituted chalcones useful as sensitizers are described in U.S. Patent No. 3,617,288. The compounds of 35 this invention can be used in photosensitive compositions in combination with other photoinitiators including the benzophenones, benzoin ethers, thioxanthone, benzil, and Michler's ketone. The compounds of this invention can be substituted for the triazines used in conjunction with dialkylamino aromatic carbonyl compounds disclosed in U.S. Patent No. 4,259,432, with 2-(benzoylmethylene)-5-benzothiazolidene thiazole-4-1 compounds disclosed in E application 0109291, May 23, 1984, with 3-keto-substituted coumarin 40 compounds disclosed in U.S. Patent No. 4,505,793, with those described in U.S. Patent No. 4,239,850, Jpn. Kokai Tokkyo Koho JP 60 60,104 (85 60104), and Ger. Offen 2,851,641.

The photoinitiators of this invention can be used with photopolymerizable compositions comprising unsaturated, free radical initiated, chain propagating addition polymerizable compound, the photoinitiator of this invention, and, optionally, one or more fillers, binders, dyes, polymerization inhibitors, color precursors, 45 oxygen scavengers, etc. The photoinitiators of this invention should be present in an amount sufficient to initiate polymerization of the polymerizable compound. For every 100 parts of polymerizable compound. there can be present from about 0.005 to about 10 parts of photoinitiator, from about 0 to about 200 parts of filler, from about 0 to about 200 parts of binder, and from about 0 to about 10 or more parts of dyes. polymerization inhibitors, color precursors, oxygen scavengers, etc., as may be needed for a particular use 50 of the photopolymerizable composition. Preferably, per each 100 parts of polymerizable compounds, there are from 1 to 7.5 parts of photoinitiator and from 25 to 150 parts of binder.

Unsaturated, free-radical initiated, chain-propagating addition polymerizable compounds suitable for the compositions of this invention include alkylene or polyalkylene glycol diacrylates, e.g., ethylene glycol diacrylate, diethylene glycol diacrylate, glycerol diacrylate, glycerol triacrylate, ethylene glycol 55 dimethacrylate, 1,3-propanediol dimethacrylate, 1,2,4-butanetriol trimethacrylate, 1,4-cyclohexanediol diacrylate, pentaerythritol tetramethacrylate, pentaerythritol triacrylate, sorbitol hexacrylate; bis[1-(3-acryloxy-2hydroxy)]-p-propoxyphenyl dimethylmethane, bis[I-(2-acryloxy)]-p-ethoxyphenyl-dimethylmethane, tris hydroxyethyl-isocyanurate trimethacrylate, the bis-acrylate and the bis-methacrylates of polyethylene

glycols of molecular weight 200-500 and the like: unsaturated amides, e.g., methylene bis-acrylamide, themselvene bis-acrylamide, themselvene bis-acrylamide, themselvene bis-acrylamide, themselvene bis-acrylamide, definyelvene triamine trisacrylamide, beta-methacrylamichetyl methacrylate; timyl esters such as divinyl succinate, drinnyl adipate, divinyl phthalate. The preferred unsaturated compounds include pentserythrich letracrylate, bis[p-q3-acryloxy-2-bydrox-propoxylphenyl] dimethylmethane, and bis[p-q3-acryloxy-phenyl] dimethylmethane. Mixtures of these esters with alkyl esters of acrylic acid and methacrylic acid, including such esters as methyl acrylate, methyl methacrylate, and pla crylate, isopropyl methacrylate, helpyl acrylate, stearyl acrylate, alkyl acrylate, and the like.

To prepare the photosensitive compositions of this invention, the components can be admixed in any order and stirred or milled to form a solution or uniform dispersion. Photosensitive elements can be made by coating the photosensitive composition on a suitable base or support and drying the coating. The dry thickness of the coating typically ranges from about 0.00005 to about 0.075 inch.

Bases or supports for the photosensithe compositions include metals, e.g., steel and aluminum plates, sheets and folis, and films or plates composed of various film-forming synthetic or high polymers, including addition polymers, e.g., vinylidane chloride, vinyl chloride, vinyl acetate, styrane, isobutylene polymers and copolymers; linear condensation polymers e.g., polyethylene tersphthalate, polyhexamethylene adipate, polyhexamethylene adipate, polyhexamethylene adipate.

The invention will be more specifically illustrated by the following examples. All values of \(\text{\lambda} max \) were measured in tetrahydrofuran, unless otherwise indicated.

Example 1

To a solution containing 0.01 mol 1-benzoyl cyclohexanol (frgacure* 184) and 12 drops din-houtytind dilaurate jn 40 ml dry toluene was added a solution of 0.01 mol 2.4-bis(trichioromethyl)-Bisozyanatc-1.3.5-triazine in toluene. The reaction mixture was stirred at room temperature under № for 24-72 hrs. The solvent was removed under reduced pressure by means of a rotary evaporator, and the residue was loaded upon a silica gel column (100 g packed with dichloromethane) and eluxed with dichloromethane. The major compound was collected and the solvent was removed by means of a rotary evaporator to afford product. The product had a melting point of 115-119 °C and a λmax of 236 nm. The structure of the product is shown below.

Example 2

The procedure of Example 1 was repeated, with the only exception being that 4-hydroxyacetophenone was used instead of 1-benzoyl cyclohexanol. The product had a melting point of 140-143 °C and a λmax of 5 280 nm, 234 nm. The structure of the product is shown below.

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The procedure of Example 1 was repeated, with the only exception being that 4-hydroxybenzophenone was used instead of 1-benzoyl cyclonexanol. The product had a melting point of 105-109 C and a kmax of 288 nm; 234 nm. The structure of the product is shown below.

Example 4

The procedure of Example 1 was repeated, with the only exception being that 4-aminobenzophenone was used instead of 1-benzoyl cyclohexanol. The product had a melting point of 225-228 °C and a λmax of 288 nm; 234 nm. The structure of the product is shown below.

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The procedure of Example 1 was repeated, with the only exception being that 2-aminoanthroquinone was used instead of 1-benzoyt cyclohexanol. The product had a mething point in exces of 280°C and a xmax of 279 nm, 241 nm. The structure of the product is shown below.

Example 6

The procedure of Example 1 was repeated, with the only exception being that 2-(hydroxymelhy)antroquince was used instead of 1-benzoyl cyclohexanol. The product had a melting point of 253-255 C and a Xmax of 255 nm. The structure of the product is shown below.

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The procedure of Exemple 1 was repeated, with the only exception being that 2-amino-9-fluorenone was used instead of 1-benzoyl cyclohexanol. The product had a melting point in excess of 260 °C and a hamax of 266 nm. The structure of the product is shown below.

Example 8

The procedure of Example 1 was repeated, with the only exception being that 4-piperazinoacetophenone was used instead of 1-benzoyl cyclohexanol. The product had a melting point of 179-182° C and a xmax of 314 mm, 238 mm. The structure of the product is shown below.

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To a solution of 1.14 g (3.3 mmol) 2,4-bis(thichromethyl)-6-isocyanato-13,5-frazine and 1.0 g (3.3 mmol) dimethyl-4-hydroxyphenylsulfonium hexafluorophosphate in 60 ml tetrahydrofuran was added 12 drops of dir-buyltin diaurate. The reaction was stirred at room temperature for 4 days and the solvent was removed on a rotary evaporator under reduced pressure. The residue was treated with hexane and the solid was filtered and dried to afford 1.01 g product. The product had a metting point of 160-163 °C and a xmax of 242 nm. The structure of the product is shown below.

Examples 10

To a solution of 1 equivalent 2-amino-4.6-bis(trichloromethyl)-1.3.5-triazine in dry toluene at 0 °C was add dropwise over a 15 minute period a solution of trichloroacelyl chloride in dry toluene. The resulting solution was heated at reflux under rititogen atmosphere for three hours and cooled to room temperature. The solvent was reduced by means of a rotary evaporator under reduced pressure. The precipitate was filtered and dried to afford product. The product had a melting point of 101-108 °C and a \text{\text{\text{max}} a \text{\text{\text{max}}} a \text{\text{\text{max}}} at 239 nm. The structure of the product is shown below.

Example 11

The procedure of Example 10 was repeated, with the only exception being that N.N-bis-(2.4-bis-(trichloromethyl)+1,3.5-friaziny+(6))-ethylenediamine was used instead of 2-amino-4.6-bis(trichloromethyl)-1,3,5-friazine. The product had a melting point of 209-213 °C and a \(\text{max} \) of 274 nm. The structure of the product is shown below.

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The procedure of Example 10 was repeated, with the only exception being that 2.6-bis-(2.4-bis-(trichloromethyl)-1,3.5-triaziny1-(6)-amino]pyridine was used instead of 2-amino-4.6-bis(trichloromethyl)-1,3.5-triazine. The product had a melting point of 243-246 °C and a λπαx of 324 nm, 233 nm. The structure of the product is shown below.

Example 13

To a solution of 2.3 mmol 2.4.8-tris(richloromethyl)-1.3.5-trisaine in 25 ml foluone was added 1 equivalent 4-piperazinoacetophenone. The reaction mixture was stirred at room temperature for 24 hours under nitrogen atmosphere. The solvent was removed by means of a rotary seporator under reduced of pressure, and the residue was dissolved in a small amount of dichloromethrane and loaded upon a column of sitica get (100 g packed in hexane) and eluted with hexane. The appropriate fractions were pooled and the solvent was removed by means of a rotary evaporator to afford product. The product had a melting point of 152-155 °C and a hamx of 312 mn, 240 nm. The structure of the product is shown below.

Example 14

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To a solution of 1.0 g (2.0 mmol) 2.4-bis(trichloromethyl)-6-[p-(2-hydroxyethoxy)shyry]-1.3.5-friazine and 0.7 f g (2.0 mmol) 2.4-bis(trichloromethyl)-6-isocyanato-1.3.5-friazine in 30 ml tetrahydrofuran was added 10 drops dibulythiofillaurate. The reaction mixture was stirred at room temperature for seventy-two hours. The solution was then concentrated on a rotary evaporator under reduced pressure. The precipitate was filtered 5 and dried to afford 1.12 g product. The product had a melting point of 172*-174* C. The structure of the product is shown below.

$$C1_{3}C = CH = CH$$

$$CC1_{3}$$

$$CC1_{3}$$

$$CC1_{3}$$

$$CC1_{3}$$

$$CC1_{3}$$

Example 15

This example illustrates the preparation of photosensitive elements using the halomethyl-1,3,5-triazines of the present invention.

A solution was prepared from 74.24 g azeotrope of 1-propanol and water (71.8% 1-propanol/28.2% water), 4.32 g pentaerythritol tetraacrylate ("Sartomer" monomer SR-295, Arco Chemical Company), 5.64 g oligomer (prepared according to U.S. Patent No. 4,228,232 and 60.9% in methyl ethyl ketone), 0.30 g triethylamine, and 14.88 g a 1:1 mixture of polyvinyl acetate-methylal resin ("Formvar" 12/85T, Union 35 Carbide Corp.) and red pigment (Pigment Red 48, C.I. 15865) (9.4% by weight solution of the azeotrope). To 2.5 g of this solution was added 2.5 mg dimethylaminobenzylacetone (DMBA), 10 mg initiator, and the resulting solution shaken in the dark for 15 minutes. The solution was filtered through glass wool and coated onto a grained, anodized aluminum plate with a #12 Mayer bar. The plate was dried at 66° C for 2 min and cooled to room temperature. To this was applied a topcoat formulation (prepared from 5.00 g carboxymethyl cellulose ether (CMC-7L), 0.26 g surfactant ("Triton" X-100 (10% in water)), and 95 g water with a #14 Mayer bar and carefully dried with a heat gun. The plates were exposed for 5 sec in air on top of a draw-down glass in a 3M Seventy exposure unit equipped with a 2 kw photopolymer bulb through a 1/2, 21 step Stouffer step tablet. The plates were soaked in the developer solution prepared from 784.40 g deionized water, 16.70 g sodium metasilicate pentahydrate, 33.40 g 1-propanol, and 0.50 g surfactant ("Dowfax-2A1", Dow Chemical Company) (45% solution in water) for 15 sec and rubbed 10 times with a 4 in. x 4 in. cotton pad. The relative sensitivities for triazines of Example 1-8, 10-12 are shown in Table 2.

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Table 2

Solid Step
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Various modifications and alterations of this invention will become apparent to those skilled in the art without departing from the scope and spirit of this invention, and it should be understood that this invention is not to be unduly limited to the illustrated embodiment set forth herein.

Claims

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- 1. A 1.3.5-triazine compound having at least one halomethyl group on a carbon atom of the triazine nucleus and having attached to another carbon atom of the triazine nucleus at least one additional photoinitiator molety, said photoinitiator molety being capable of initiating free radical or ionic chain polymerization upon exposure to actinic radiation.
 - 2. The compound of claim 1 wherein said at least one halomethyl group is a trihalomethyl group.
- 3. The compound of claim 2 wherein said trihalomethyl group is a member selected from the group consisting of trichloromethyl group, tribromomethyl group, and trilodomethyl group.
- 4. The compound of claim 1 wherein said photoinitiator molety is selected from the group consisting of benzoin group, benzoin alkyl ether group, acetophenone group, dialkoxyacetophenone group, benzophenone group, fluorenone group, anthraquinone group, thioxanthone group, triary/sulfonium group, diaryliodonium group, acytoxime group, azide group, diazonium group, 3-ketocoumarin group. bisimidazole group, and halomethyl-1,3,5-triazine group.
- 5. A radiation-sensitive composition comprising: (1) an ethylenically unsaturated, polymerizable compound, and (2) a compound according to claim 1.
 - 6. A compound having the formula:

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A represents a member selected from the group consisting of mono-, di- and trihalomethyl groups, 55 Y represents a member selected from the group consisting of A, L-P, NH2, NH2, NH2, OR, and R, where R independently represents a substituted or unsubstituted alkyl group, or a substituted or unsubstituted aryl group, R represents a substituted or unsubstituted alkyl group, a substituted or unsubstituted aryl group, a substituted or unsubstituted alkenyl group, or a substituted or unsubstituted heterocyclic aromatic group,

P represents a photoinitiator moiety capable of initiating free radical or ionic chain polymerization upon exposure to actinic radiation, and

L represents a group or a covalent bond linking the photoinitiator moiety to the triazine nucleus.

- 7. The compound of claim 6 wherein A represents a trihalomethyl group.
- 8. The compound of claim 7 wherein the trihalomethyl group is a member selected from the group consisting of trichloromethyl, tribromomethyl, triidomethyl.
- The compound of claim 8 wherein the trihalomethyl group is a member selected from the group consisting of trichloromethyl and tribromomethyl.
- 10. The compound of claim 6 wherein the photoinitiator molety is selected from the group consisting of benzoin group, benzoin alkey letter group, acetophenone group, dialkovszectophenone group, thiorenone group, thiorenone group, thiorenone group, thiorenone group, acetyloxime group, acetyloxim
 - 11. The compound of claim 6 wherein Y represents A.
 - 12. The compound of claim 6 wherein Y represents L-P.
 - 13. The compound of claim 6 wherein R represents a substituted or unsubstituted aryl group.
 - 14. The compound of claim 6 wherein R represents a substituted or unsubstituted heterocyclic aromatic group.
 - 15. The compound of claim 6 wherein R represents a substituted or unsubstituted alkenyl group.
 - 16. A radiation-sensitive composition comprising: (1) an ethylenically unsaturated, polymerizable compound, and (2) a compound according to claim 6.

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Category	Citation of document with in of relevant pas		Relevant te claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.5)
Х	EP-A-0 305 115 (VIC * Whole document, es	KERS PLC) specially pages 2,7	1-16	G 03 F 7/029 C 07 D 251/22 C 07 D 401/14
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				G 03 F 7/00
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	The present search report has b	een drawn up for all claims		
TH	Place of search HAGUE	Date of completion of the search		Examiner M. M. B.
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- Y: particularly relevant if combined document of the same category A: technological background O: non-written disclosure P: intermediate document

- D: document cited in the application L: document cited for other reasons
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